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Bezeichnung der Erfindung/Title of the invention/Titre de l'invention:
(Falls die Bezeichnung der Erfindung nicht angegeben ist, siehe Beschreibung.
If no title is shown please refer to the description.
Si aucun titre n'est indiqué se referer à la description.)

Microstructures

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MicrostructuresField of the Invention

5 The present invention generally relate to microstructures and particularly relates to microstructures incorporating carbon nanotube tips for applications including microscopy and data storage.

10

Background of the Invention

15 Atomic Force Microscopy (AFM) is a well known form of Scanning Probe Microscopy (SPM). AFM is conventionally employed in range of surface imaging, nanometer scale analysis and manipulation applications. Imaging is achieved via AFM by detecting forces occurring between a sensor in form of a tip at one end of a cantilever and a sample to be imaged. The tip and cantilever in combination will be herein after referred to

20 as a tip assembly in the interests of convenience. The tip is a relatively reliable tool for providing highly localized confinement of interaction. This property has opened up a range of applications in the nanotechnology field. For example, in "The "Millipede" - More than one thousand tips for

25 future AFM data storage", P. Vettiger et al, IBM Journal of Research and Development. Vol. 44 No. 3, May 2000, there is described a data storage device based on AFM probe technology providing smaller form factor, higher capacity, lower power, and lower cost than conventional memory devices. The storage

30 density achievable in such a storage device depends on the durability and quality of the tip. It would therefore be desirable to provide such tips with optimal durability and quality via a cost effective process. It would also be desirable to produce highly integrated arrays of such sensors.

In a conventional AFM instrument, the tip assembly, which is responsible for the spatial resolution of the instrument, contributes a significant portion of the total cost of ownership (TCO) of the instrument. The cantilever of the tip assembly is typically hand-fabricated or batch microfabricated cantilever. In situ within the instrument, one end of the cantilever is fixed and the other remains free. The tip is located at the free end. In operation, the cantilever permits sensing of a force interaction between the tip and the surface of the probed sample. A surface observation made by such an instrument is a function of the sample surface topography and the shape of the tip. To minimize noise, it is desirable not only to make the tip as sharp as possible, but also to make the aspect ratio of the tip as high as possible. The aspect ratio of the tip is an inverse function of the cone angle of the tip. However, as the aspect ratio of a tip is increased, so the tip becomes more fragile and subject to wear. Usually, the shape of the tip is determined by way of tradeoff between the tip robustness and the imaging quality.

An example of a conventional tip assembly comprises a microfabricated cantilever with an integrated silicon tip. While allowing for some economy of scale in production costs over hand-fabricated cantilevers, this technology is nonetheless still relatively expensive. In addition, the silicon tip is subject to wear during normal operation. Wearing of the tip can lead to inconsistent imaging results. Wearing of the tip also makes the tip a consumable of the instrument, introducing a need for regular tip monitoring and replacement time costs. It would be desirable to reduce such monitoring and tip replacement requirements.

WO 99/56176 discloses a method of manufacturing a tipped cantilever comprising forming a tip-like indent in a substrate, depositing a photoresist layer which fills the

tip-like indent and covers at least a part of said substrate, and photolithographically structuring the photoresist layer to form the tipped cantilever with tip out of the photoresist. This technique lends itself well to batch microfabrication of

5 plastic cantilevers at a lower cost than the aforementioned silicon technology.

In Review of Scientific Instruments, vol. 70, no. 5, May 1999, pages 2398-2401, G. Genolet et al., "Soft, entirely

10 photoplastic probes for scanning force microscopy", there is disclosed a probe made entirely of plastic material for scanning probe microscopy. A polymer is used for forming the cantilever. The polymer provides mechanical properties that are difficult to achieve with classical silicon technology.

15 The fabrication process is a batch process in which the integrated tip and the lever are defined in one photolithography step.

Applied Physical Letters, vol. 77, no. 21, 20 November 2000,

20 R. Stevens et al., "Improved fabrication approach for carbon nanotube probe devices", discloses an improved process for simple and efficient fabrication of carbon nanotube probe devices. The process requires two steps. First, a nanotube cartridge is created using chemical vapor deposition. Then,

25 the nanotubes are transferred from the cartridge to a device using an electric field.

Carbon nanotube mounted Si and/or silicon nitride tips are commercially available, for example, from PIEZOMAX

30 Technologies Inc. The nanotube is grown and then mounted on such tips manually. The length of the attached nanotube is then iteratively tuned via a series of alternating shortening and re-inspection steps. It will be appreciated that this serial production process is both time consuming, expensive,

35 and unsuitable for batch fabrication. Additionally, the

robustness of the tip assembly is governed by the bond between the nanotube and the tip.

5 In general, the quality and durability of tip assemblies produced according to the aforementioned conventional techniques present limitations is less than optimal for the range of applications now envisaged for such instruments. Furthermore, the reproducibility of such tip assemblies via conventional methods is difficult to achieve without incurring 10 additional cost. It would desirable therefore, as mentioned earlier, to provide microstructures such as tip assemblies of optimal quality, durability and versatility that can be manufactured cheaply.

15 Summary of the Invention

In accordance with the present invention, there is now provided a method for forming a microstructure, comprising: 20 depositing a seed material on a substrate; growing a nanotube from the seed material; depositing microstructure material on the substrate to embed the nanotube in the microstructure material; and, detaching the substrate to release the microstructure.

25 The method preferably further comprises shaping the microstructure material prior to release of the microstructure. The microstructure material may comprise any one of a polymer, a dielectric material, a metal, and polysilicon. The substrate may be formed from any one of 30 Silicon, glass, quartz, ceramics, and plastic. The seed material preferably comprises alternating layers of a first and second precursor materials. The first precursor material preferably comprises a fullerene material, and the second precursor material preferably comprises field sensitive 35 material. The fullerene material may comprise any one of

Carbon 60 and Carbon 82, and the field sensitive material may comprise any one of Ni, Co, Fe and Mo. In a preferred embodiment of the present invention, the fullerene material comprises Carbon 60, and the field sensitive material

5 comprises Nickel. The growing of the nanotube preferably comprises: heating the substrate in vacuum conditions; and applying a field to the substrate. The vacuum conditions, in a preferred embodiment of the present invention, comprise oxygen pressure greater than 10^{-5} mbar. The heating preferably

10 comprises elevating the temperature of the substrate to between 900 and to 1000 degrees Centrigade. The applied field may comprise a magnetic field. The magnetic field is preferably applied orthogonally to the surface of the substrate. In preferred embodiments of the present invention,

15 the magnetic field is greater than or equal to 50 Gauss. Additionally or alternatively, the applied field may comprise a electric field. The electric field is preferably applied orthogonally to the surface of the substrate. The detaching preferably comprises depositing a sacrificial layer on the

20 surface of the substrate prior to deposition of the cantilever material; and immersing the sacrificial layer in an electrolyte after deposition of the microstructure material. The sacrificial layer preferably comprises an anode sub-layer and a cathode sub-layer. The anode sub-layer may comprise any

25 one of Al, Zn, Cr, Fe, and Co, and the cathode sub-layer comprises a noble metal. The cathode sub-layer may comprise any of Au, Pd, Pt, Ag, and Cu.

30 In a preferred embodiment of the present invention, the depositing of the seed material comprises: depositing a photo-resist layer on the substrate; forming an aperture in the resist layer; masking the substrate with a resist layer to locate the seed material at a site on the substrate defined by the aperture; and, removing the resist layer to remove surplus seed material. The forming of the aperture preferably

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comprises under-etching the resist layer to produce a cavity in the resist layer.

Another embodiment of the present invention comprises forming
5 a tip image in a substrate to produce a mold for receiving the
microstructure material. The forming of the tip image
preferably comprises: depositing a photo-resist layer on the
substrate; forming an aperture in the photo-resist layer; and,
10 under etching the substrate beneath the photo-resist layer to
create the tip image. The depositing of the seed material
preferably comprises: masking the substrate with the resist
layer to locate the seed material in the apex of the tip
image; and, removing the resist layer to remove surplus seed
material.

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Viewing the present invention from another aspect, there is provided a microstructure comprising a body portion and a nanotube embedded in the body portion.

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The body portion may comprise a cantilever beam with the nanotube embedded in and extending laterally from one end of the beam. Equally or additionally, the body portion may comprise a laterally extending tip with the nanotube embedded
25 in and extending from the apex of the tip. The body portion may comprise any one of a polymer, a dielectric material, a metal, and polysilicon.

It will be appreciated that the present invention extends to a
30 micromechanical sensor comprising a microstructure as hereinbefore described. The present invention also extends to a data storage device comprising an array of such sensors for reading or writing data onto a storage surface.

35 The present invention advantageously provides a much stronger

tip assembly than the conventional tip assemblies herein before described because the nanotube is embedded in the tip material. Another advantage is that the nanotube provides a major improvement in imaging resolution because it offers a 5 smaller apex radius to the surface under analysis. In addition, the nanotube permits imaging of rougher surfaces because it provides an much higher height to width aspect ratio. Furthermore, the nanotube has an electrical resilience, thereby imparting additional robustness and improved contact 10 force to the tip assembly. Still furthermore, a nanotube, as a macromolecule may also provide a predictable platform for tip functionalization by other molecules, such as those of a biochemical nature, allowing the production of scanning probe 15 chemical and biological sensors.

15 Preferably, the sensor comprises a bundle of nanotubes extending from the apex portion of the tip. The bundle preferably protrudes from the apex portion of the tip. The cantilever beam may comprise any one of photoplastic material, 20 metal, and polysilicon.

25 The present invention also extends to a data storage device comprising an array of sensors as hereinbefore described for reading or writing data onto a storage surface.

Other advantages of the present invention may be apparent to those skilled in the art based on a perusal of the following 30 description of preferred embodiments.

30 Brief Description of the Drawings

Preferred embodiments of the present invention will now be described, by way of example only, with reference to the accompanying drawings, in which:

Figures 1A to 1J depict steps of a method of fabricating a microstructure according to the present invention;

5 Figures 2A to 2I depict steps of another method of fabricating a microstructure according to the present invention; and, Figure 3 is a block diagram of a local probe storage device including an array of tips.

10 Detailed Description of Preferred Embodiments

An example of a method for fabricating a microstructure according to the present invention will now be described with reference to Figure 1. The Figure 1 method will be described with reference to fabrication of a microstructure in the form 15 of a micromechanical cantilever tip assembly having an carbon nanotube crystal embedded in a tip formed from a moldable material. However, it will be appreciated that the method is equally applicable to fabrication of other microstructures having nanotubes embedded therein.

20 Referring first to Figure 1A, a layer 2 of a photoresist material is initially deposited on a substrate 1. In a preferred embodiment of the present invention, the substrate 1 is formed of Silicon. However, in other embodiments of the 25 present invention, the substrate 1 may be formed of a different material, such as glass, quartz, ceramics, plastic, and the like. Other suitable materials will be apparent to those skilled in the art. The photo resist layer 2 is deposited, in a particularly preferred embodiment of the 30 present invention, by spin coating. However, in other embodiments of the present invention, the resist layer 2 may be deposited by a different technique, such as physical vapor deposition or chemical vapor deposition. An aperture 3 of about 300 nm in diameter is opened in the resist layer 2 to 35 expose the underlying Silicon of the substrate 1. In a

preferred embodiment of the present invention, the aperture 3 is opened by photolithography. However, it will be appreciated that, in other embodiments of the present invention, different techniques may be employed for opening the aperture 3.

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Referring now to Figure 1B, a conical cavity 4 is then formed in substrate 1 beneath the resist layer 2. The cavity 4 is substantially the image of the tip of the sensor to be produced. In a particularly preferred embodiment of the 10 present invention, the cavity 4 is formed by under-etching the substrate 1 beneath the resist layer 2. Isotropic or anisotropic plasma etching may be employed to provide such under etching. Different techniques for forming the cavity 4 may be employed in other embodiments of the present invention.

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Referring to Figure 1C, a nanotube seed layer 5 is then deposited on the resist layer 2. In a particularly preferred embodiment of the present invention, the seed layer 5 is 20 deposited by thermal evaporation. However, in other embodiments of the present invention, the seed layer 5 may be deposited by a different technique, such as physical vapor deposition or chemical vapor deposition.

25 In a preferred embodiment of the present invention in which the substrate 1 is formed from Silicon, a protective layer is disposed at least at the apex of the cavity 4. The protective layer prevents diffusion of materials constituting the seed layer 5 into the substrate 1. In a particularly preferred 30 embodiment of the present invention, the protective layer is formed from Silicon Dioxide. In another preferred embodiment of the present invention, the protective layer is formed from Molybdenum.

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The seed layer 5 comprises a first precursor material comprising fullerene molecules such C60, C70, C82, or C116, and a second precursor material comprising a catalyst. In particularly preferred embodiments of the present invention, 5 the first precursor materials comprises fullerene molecule C60 and the catalyst comprises a magnetic material such as one or more metals selected from the group Ni, Co, Fe, Mo.

In preferred embodiments of the present invention, the first 10 and second precursor materials are deposited in alternating layers. Preferably, between five and ten layers of each precursor material are interlaced in this manner. In a particularly preferred embodiment of the present invention, each layer of precursor material has a thickness of between 5 15 and 30 nm. During deposition of the seed layer 5 by evaporation, the resist layer 2 acts as a shadow mask directing a portion of the seed layer 5 in the form of a pellet 6 to the apex of the cavity 4. Conditions for controlled nanotube growth such as seed material localization 20 and presence of a micron scale defect are thus optimized in the apex of the cavity 4.

The evaporation of the precursor materials may be performed at 25 pressure of around 10^{-9} Torr. Control of such evaporation can be achieved via an electromechanical shutter with an in situ balance for monitoring the deposition rate of the precursor materials. The evaporation can be thus regulated such that the thickness of the layers decreases with their distance from the substrate 1. This decreasing thickness increases the yield. 30 Also, the reduction in thickness may also lead to less of the catalyst being transported towards the tip of the growing nanotube crystal. The evaporation of some desirable catalysts, such as Nickel for example, is technically difficult. This difficulty makes it desirable to utilize only the minimum 35 necessary amount for the manufacturing process. Hence the

amount of catalyst material can be reduced by the thinner layers. Because the growth of the crystal begins at the base of the pellet 6, less material transport from the layers remote from the substrate 1 occurs with the layers of reduced thickness.

Referring to Figure 1D, the resist layer 2 is then removed thereby lifting-off the bulk of the seed layer 5. However, the pellet 6 remains in the apex of the cavity 4. In a particularly preferred embodiment of the present invention, the resist layer 2 is removed by wet or dry etching. Different techniques for removing the resist layer 2 may be employed in other embodiments of the present invention. In preferred embodiments of the present invention, the pellet 6 has a diameter of the order of 300 nm. However in other embodiments of the present invention, the seed pellet 6 may have a larger or smaller diameter.

With reference to Figure 1E, a nanotube crystal 7 is then grown from the pellet 6 at the apex of the cavity 4. The crystal 7 may comprises a bundle of nanotubes or, more preferably, a single nanotube. Growth of the crystal 7 is activated, in preferred embodiments of the present invention, by heating the substrate 1 in vacuum conditions and in the presence of a magnetic or electric field or in combination of magnetic and electric fields. In some embodiments of the present invention, the substrate 1 may be heated in substantially an inert gas atmosphere.

In preferred embodiments of the present invention, the substrate 1 is heated to between 900 and 950 degrees centigrade. The vacuum conditions, in a preferred embodiment of the present invention are provided by a oxygen pressure greater than 10^{-6} mbar. In a preferred embodiment of the present invention, the heating is performed in the presence of

a magnetic field in the region of 50 Gauss or higher orthogonal to the plane of the substrate 1. The crystal 7 grows in the direction of the applied magnetic field, thus extending in a direction normal to the plane of the substrate 5 1.

In another embodiment of the present invention, growth of the crystal 7 is performed in the presence of an electric field extending perpendicular to the plane of substrate 1 to promote 10 growth of the crystal 7 in a direction perpendicular to the plane of the substrate 1.

In a particularly preferred embodiment of the present invention, growth of the crystal 7 is stimulated by heating 15 the substrate to 950 degrees centigrade in a vacuum of 10^{-6} Torr and in the presence of a perpendicular magnetic field for a period of between around three minutes and around 30 minutes.

20 Generally, adequate growth can be obtained with a heating time in the minute range. Longer heating does not significantly improve the result. In another preferred embodiment of the present invention, a vacuum of 10^{-9} Torr is used.

25 The dimensions of the pellet 6 determine the total amount of the precursor materials that are involved in the growth of the nanotube crystal 7. The pellet 6 thus provides a reservoir of precursor materials from which the nanotube crystal material is supplied. The predetermination of the material supply in 30 the pellet 6 has the effect that the different precursor materials used in the growth of the nanotube crystal 7 are predetermined in terms of amount and position. The movement of the molecules of the precursor materials is hence confined within the pellet 6, thereby leading to more defined growth 35 and location. The relative concentrations of the precursor

materials can affect growth of nanotube crystal 7. Again however, the confinement of the precursor materials in the pellet 6 leads to a more defined ratio between the two precursor materials.

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The stronger the applied field, the larger the pellet 6 that can be grown into the crystal 7 because the force that directs the second precursor material is stronger. The shape of the pellet 6 need not be round or square. Other shapes are 10 possible. However, for symmetry reasons a circular shape is preferred. The bundle may range from one to several million nanotubes.

15 A nucleation site may be placed on the substrate 1 to enable controlled positioning of the nanotube crystal 7. The nucleation site may be created for example by evaporating through the resist layer 2 a material such as tungsten. Because the resist layer 2 has a shadowing effect, an evaporator for the nucleation material, situated sufficiently 20 apart from the evaporators for the precursor materials can automatically generate the nucleation site near the pellet 6. It is desirable to locate the evaporators for the precursor materials as close together as possible to avoid lateral misalignment of the alternate precursor layers in the pellet 25 6.

30 As indicated earlier, electromechanical shuttering combined with an in situ quartz crystal micro-balance to monitor deposition rates, may be used to ensure that both fullerene and catalyst materials are evaporated sequentially to produce the desired structure of the pellet 6. This technique produces the pellet 6 at a site determined by the relative position of the aperture 3 and the substrate 1. The choice of substrate 1 35 can be influenced by the choice of fullerene and catalyst materials.

Acceptable results can be achieved with the substrate 1 being made of Silicon Dioxide or molybdenum in the form of a solid film sputtered on to a Silicon wafer.

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In a particularly preferred embodiment of the present invention, after evaporation of a pellet 6 of alternating layers Carbon 60 and Nickel on the substrate 1, the arrangement is heated to 950°C in a vacuum of 10^{-6} mbar for a 10 time which is chosen to lie between a few minutes and an hour. Growth of the resulting nanotubes 7 is oriented normal to the surface of the substrate 1 by immersing the substrate in a magnetic field oriented perpendicular to the substrate surface during heat treatment. A field strength of around 1.5 Tesla 15 achieves desirable results. The application of an electric, electromagnetic or magnetic field directs the self-assembly and organization of the single-walled carbon nanotubes. The electric field can be an AC or DC field. The electromagnetic field can also be an optical near field such as that of a 20 laser. Also, atomic forces or Van der Waals forces may be applied during growth.

The direction of movement of the second precursor material is governed by the applied field. Hence the field and the second 25 precursor material cooperate in that the second precursor material is moveable by the field. Such a property can be that the second precursor material is magnetic or bears an electric charge or a combination thereof, or has some other 30 property that is influenced by the applied field to exert a moving force.

The nanotube bundle 7 produced in preferred embodiments of the present invention typically has a diameter between 40nm and 900nm with length up to 2 microns. The nanotubes therein are 35 straight are single-wall carbon nanotubes forming straight

5 bundles normal to the substrate. The wall diameters in the bundle are substantially uniform and range from about 1.4nm to 2.3nm. There is an inverse correlation between wall and bundle diameter in that small wall diameters are predominant in large diameter bundles whereas large wall diameters are prevalent in small diameter bundles. A typical bundle as a length of about 750 nm and a diameter of between around 20 nm and around 100 nm with a curved end cap, with each nanotube therein having a diameter of around 1.6 nm. The nanotubes are substantially 10 free of inhomogeneity or defect. This remarkable structural perfection is a characteristic of nanotubes produced using the described nanotube growth technique. Further details of this technique are provided in our co-pending patent application reference CH9-2001-0071.

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20 Turning now to Figure 1F, a sacrificial layer 8 is now deposited on the substrate 1. The sacrificial layer 8 may be deposited by any of a range of techniques, including physical vapor deposition or chemical vapor deposition.

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25 The sacrificial layer 8 comprises two electrically conductive electrode materials having different oxido-reduction potentials. The deposition of the two materials constituting the sacrificial layer 8 is performed in such a way that an electrical contact is present between them. The materials 30 cooperate to form a galvanic cell, with each material forming a different electrode of the cell. The electrode with the higher oxido-reduction potential forms the cathode and the electrode having the lower oxido-reduction potential forms the anode. The cell is of sufficiently high electrochemical potential that one of the materials is electro-etched away when the sacrificial layer 8 is immersed in a suitable electrolyte. In a preferred embodiment of the present invention, the cathode is deposited first, and the anode is 35 subsequently deposited on top of the anode so that the cathode

is not released with the sensor to be produced. The electrodes may be formed from a metal or doped semiconductor. The cathode may be provided, in some embodiments of the present invention by the substrate 1.

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The electrodes may be formed from metals. Preferably, the cathode comprises a noble metal such as Au, Pd, Pt, Ag, Cu or the like and the anode comprises a metal selected from a group comprising Al, Zn, Cr, Fe, Co or the like, or a doped semiconductor.

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The difference between the oxido-reduction potentials of the anode and the cathode should be as large as possible to achieve the maximum electro-etching efficiency. The

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electrolyte may be introduced in the form of either a liquid or a vapour environment to form a galvanic cell with an electrical potential high enough to etch or enhance the etching of the anode. In a particularly preferred embodiment of the present invention, the electrolyte used is an acidic solution known to etch the anode material. The anode and cathode can be very thin (in the range of about 10 nm) yet nevertheless produce a highly controllable and fast under etching.

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Further details of this sacrificial layer release technique are provided in our co-pending patent application reference CH9-2000-0081. An advantage of employing the sacrificial layer release technique hereinbefore described for release of microstructures from moulds is that the technique is well suited to releasing microstructures from pre-structured substrates.

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With reference to Figure 1G, a layer of cantilever material 9 is then deposited onto the sacrificial layer 8. The cantilever material 9 may be deposited by any of a range of techniques,

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including physical vapor deposition or chemical vapor deposition. Referring to Figure 1H, the cantilever material is then shaped. Turning to Figure 1I, a body portion 10 of the cantilever is then deposited on the cantilever layer 9. The 5 body portion is then shaped. The shaping steps hereinbefore described with reference to Figures 1H and 1I may be implemented by any of a range of techniques, including lithography, wet etching, and plasma etching.

10 In preferred embodiments of the present invention, the body portion 10 and the cantilever layer 9 are formed from the same cantilever material. That cantilever material may comprise one or more of a range of materials, such as polymers, dielectric materials, metals, or polysilicon. In particularly preferred 15 embodiments of the present invention, the cantilever material comprises a photo-structurable polymer such as SU-8.

Referring to now Figure 1J, the completed cantilever 11 is then released from the substrate 1 by immersing the substrate 20 1 in an electrolyte environment and electro-etching the sacrificial layer 8 as herein before described. This release process exposes an end of the nanotube crystal 7 and leaves the cantilever layer 9 and body 10 unaffected. The thickness of the sacrificial layer 8 determines the length and thus 25 mechanical properties of the nanotube crystal 7. The lateral mechanical rigidity of the nanotube crystal 7 depends on its length and the aspect ratio.

In a preferred embodiment of the present invention 30 hereinbefore described, a sacrificial layer 8 was employed in the release of the cantilever 11. However, in other embodiments of the present invention, other release techniques may be employed. For example, release may also be achieved simply having a adhesion lost while drawing the cantilever 11 35 out of the substrate 11. The length of the nanotube crystal 7

may be tuned by the thickness of the sacrificial layer 8. Alternatively however, the length of the nanotube crystal 7 may be tuned by depositing a buffer layer (not shown) between the sacrificial layer 8 and the cantilever material 9. The 5 buffer layer is then removed after lift off. In a particularly preferred embodiment of the present invention, the buffer layer is removed by etching, with the cantilever 11 remaining unaffected by the etchant. In another preferred embodiment of the present invention, the buffer layer is omitted and, after 10 liftoff, the cantilever is deliberately thinned down by etchant to expose the embedded nanotube crystal 7.

Another example of a method for fabricating a microstructure according to the present invention will now be described with 15 reference to Figure 2. The Figure 2 method is variation of the method hereinbefore described with reference to Figure 1. In addition, the Figure 2 method will be described with reference to fabrication of a microstructure again in the form of a micromechanical cantilever tip assembly having an carbon 20 nanotube crystal embedded in a moldable material. However, it will also be appreciated that the Figure 2 method is equally applicable to fabrication of other microstructures having nanotubes embedded therein.

25 Referring to Figure 2A, the photo-resist layer 2 is again deposited on the substrate 2.

Turning to Figure 2B, the aperture 3 is again formed in the resist layer 2 by, for example, exposure and development of 30 the resist layer 2. However, the aperture 3 is now under-etched to form a cavity in the resist layer 2 having a necked opening.

Referring to Figure 2C, the pellet 6 is now deposited on the 35 substrate 1 via the necked opening in the resist layer 2 using

the shadow masking technique hereinbefore described with reference to Figure 1.

5 Turning to Figure 2D, the resist layer 2 is then removed from the substrate 2 leaving the pellet 6 remaining on the substrate 2.

10 With reference to Figure 2E, the nanotube crystal 7 is then grown according to the technique hereinbefore described with reference to Figure 1.

15 Referring to Figure 2F, a layer of cantilever material 9 is then deposited on the substrate 1, engulfing the nanotube crystal 7. The cantilever material 9 is preferably deposited by spin coating the substrate 1. In a particular preferred embodiment of the present invention, the cantilever material is a photo-structurable polymer such as SU-8.

20 Referring to Figure 2G, the cantilever material 9 is then structured in situ on the substrate 1.

25 Turning now to Figure 2H, the cantilever material 9, together with the nanotube embedded therein, is released from the substrate 1. The release of the cantilever material 9 may be effected, for example, by one of the release techniques hereinbefore described with reference to Figure 1. In the event that sacrificial layer release technique hereinbefore described with reference to Figure 1 is employed, the sacrificial layer is preferably deposited on the substrate 1 prior to deposition of the resist layer 2 referred to in connection with Figure 2A.

30 35 With reference to Figure 2I, the face of the cantilever material previously in contact with the substrate 1 is now thinned to expose an end of the nanotube crystal 7. The other

end of the crystal 78 remains embedded in the cantilever material 9.

With reference to Figure 3, a cantilever manufactured
5 according to a method as hereinbefore described may find application in a two dimensional cantilever sensor array 20 for a local probe data storage device 21. In such a device 21, the sensor array 20 is scanned relative to a storage surface 22 in orthogonal directions parallel to the surface 22.
10 Indentations formed in the surface 22 and representative of recorded data are detected by the tips 23 in the array 20 as they are scanned across the surface 22. Equally, the tips 23 can be energized to form indentations in the surface 22 thereby recording data in the surface 22. Further details of
15 such a storage device 21 are provided in the aforementioned Vettiger et al. article. Cantilever sensors produced in accordance with the present invention are especially useful in such storage devices because, having a tip of only few tens of nano-meters in diameter, such tips provide a much higher
20 aspect ratio than conventional tips. The indent dimension made in the polymer will not be wider when using a thicker polymer film (thick polymer film means better read back sensitivity). As for imaging wear of the nanotube will not affect the resolution, leading to the storage density. Also, the thermal
25 conductivity of the nanotube crystal is preferable to that of a silicon tip.

Preferred embodiments of the present invention have been hereinbefore described with reference to microstructures for
30 microscopy and data storage applications. However, it will be appreciated that the present invention is equally applicable to microstructures for a range of other applications. For example, the present invention is equally applicable to charge emissive microstructures such a field emission tips. Such tips
35 may be employed in a range of different application, including display devices.

CLAIMS

5 1. Method for forming a microstructure, comprising:
depositing a seed material on a substrate; growing a
nanotube from the seed material; depositing
microstructure material on the substrate to embed the
nanotube in the microstructure material; and, detaching
10 the substrate to release the microstructure.

15 2. Method as claimed in claim 1, comprising shaping the
microstructure material prior to release of the
microstructure.

3. Method as claimed in claim 1 or claim 2, wherein the
microstructure material comprises any one of a polymer, a
dielectric material, a metal, and polysilicon.

20 4. Method as claimed in any preceding, wherein the substrate
is formed from any one of Silicon, glass, quartz,
ceramics, and plastic.

25 5. Method as claimed in any preceding claim, wherein the
seed material comprises alternating layers of a first and
second precursor materials.

30 6. Method as claimed in claim 5, wherein the first precursor
material comprises a fullerene material, and the second
precursor material comprises field sensitive material.

35 7. Method as claimed in claim 6, wherein the fullerene
material comprises any one of Carbon 60 and Carbon 82,
and the field sensitive material comprises any one of Ni,
Co, Fe and Mo.

8. Method as claimed in claim 7. Wherein the fullerene material comprises Carbon 60, and the field sensitive material comprises Nickel.

5

9. Method as claimed in any preceding claim, wherein the growing of the nanotube comprises: heating the substrate in vacuum conditions; and applying a field to the substrate.

10

10. Method as claimed in claim 9, wherein the vacuum conditions comprise oxygen pressure greater than 10^{-5} mbar.

15

11. Method as claimed in claim 9 or claim 10, wherein the heating comprises elevating the temperature of the substrate to between 900 and to 1000 degrees Centrigade.

20

12. Method as claimed in any of claims 9 to 11, wherein the applied field comprises a magnetic field.

25

13. Method as claimed in claims 12, wherein the magnetic field is applied orthogonally to the surface of the substrate.

14. Method as claimed in claim 13, wherein the magnetic field is greater than or equal to 50 Gauss.

15

15. Method as claimed in any of claims 9 to 11, wherein the applied field comprises a electric field.

10

16. Method as claimed in claims 14, wherein the electric field is applied orthogonally to the surface of the substrate.

15

17. Method as claimed in any preceding claim, wherein the
detaching comprises depositing a sacrificial layer on the
surface of the substrate prior to deposition of the
cantilever material; and immersing the sacrificial layer
in an electrolyte after deposition of the microstructure
material.

5

18. Method as claimed in claim 17, wherein the sacrificial
layer comprises an anode sub-layer and a cathode
sub-layer.

10

19. Method as claimed in claim 18, wherein the anode
sub-layer comprises any one of Al, Zn, Cr, Fe, and Co,
15 and the cathode sub-layer comprises a noble metal.

20. Method as claimed in claim 19, wherein the cathode
sub-layer comprises any of Au, Pd, Pt, Ag, and Cu.

20 22. Method as claimed in any preceding claim, wherein the
depositing of the seed material comprises: depositing a
photo-resist layer on the substrate; forming an aperture
in the resist layer; masking the substrate with a resist
layer to locate the seed material at a site on the
substrate defined by the aperture; and, removing the
25 resist layer to remove surplus seed material.

23. Method as claimed in claim 22, wherein the forming of the
aperture comprises under-etching the resist layer to
30 produce a cavity in the resist layer.

24. Method as claimed in any claim preceding claim 22,
comprising forming a tip image in a substrate to produce
a mold for receiving the microstructure material.

35

25. Method as claimed in claim 24, wherein the forming of the tip image comprises: depositing a photo-resist layer on the substrate; forming an aperture in the photo-resist layer; and, under etching the substrate beneath the photo-resist layer to create the tip image.

5

26. Method as claimed in claim 25, wherein the depositing of the seed material comprises: masking the substrate with the resist layer to locate the seed material in the apex of the tip image; and, removing the resist layer to remove surplus seed material.

10

27. A microstructure comprising a body portion and a nanotube embedded in the body portion.

15

28. A microstructure as claimed in claim 27, wherein the body portion comprises a cantilever beam and the nanotube is embedded in and extends laterally from one end of the beam.

20

29. A microstructure as claimed in claim 27 or claim 28, wherein the body portion comprises a laterally extending tip and the nanotube is embedded in and extends from the apex of the tip.

25

30. A microstructure as claimed in any of claims 27 to 29, wherein the body portion comprises any one of a polymer, a dielectric material, a metal, and polysilicon.

30

31. A micromechanical sensor comprising a microstructure as claimed in any of claims 27 to 30.

35

32. A data storage device comprising an array of sensors as claimed in claim 31 for reading or writing data onto a storage surface.

ABSTRACT

A method for forming a microstructure is described. The method comprises: depositing a seed material on a substrate; growing a nanotube from the seed material; depositing microstructure material on the substrate to embed the nanotube in the microstructure material; and, detaching the substrate to release the microstructure. The resulting microstructure comprises a body portion and a nanotube embedded in the body portion.

CA9-2001-0024

1/6

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FIG. 1A

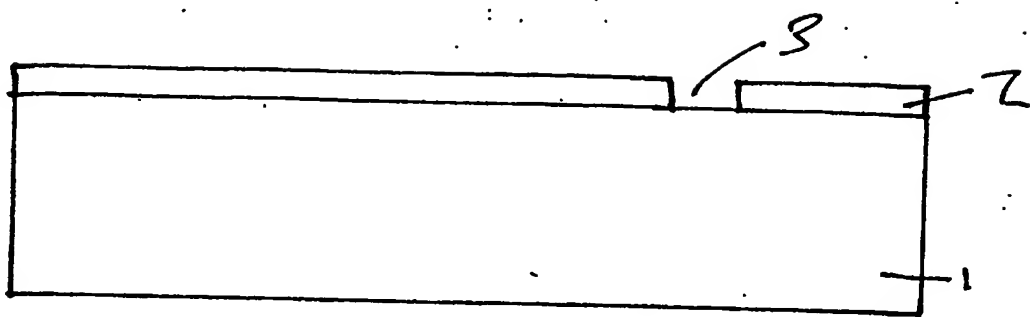


FIG. 1B

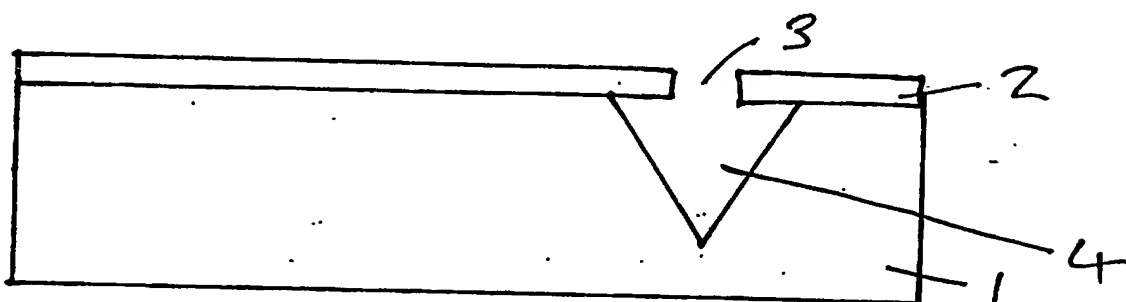


FIG. 1C

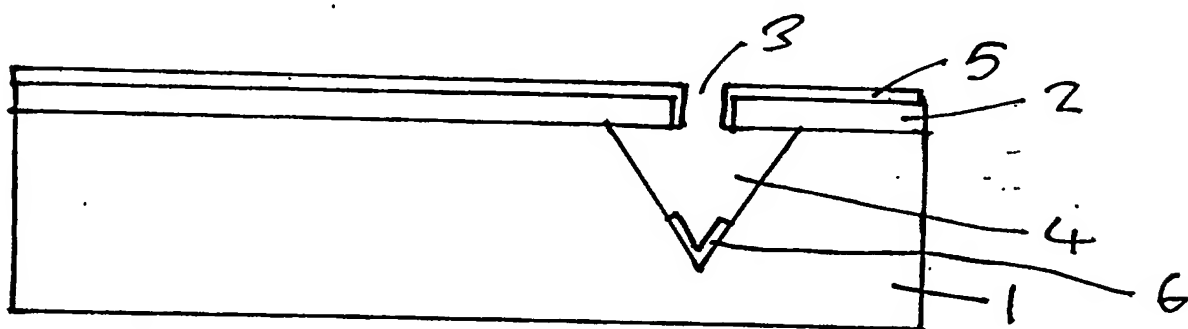


FIG. 1D

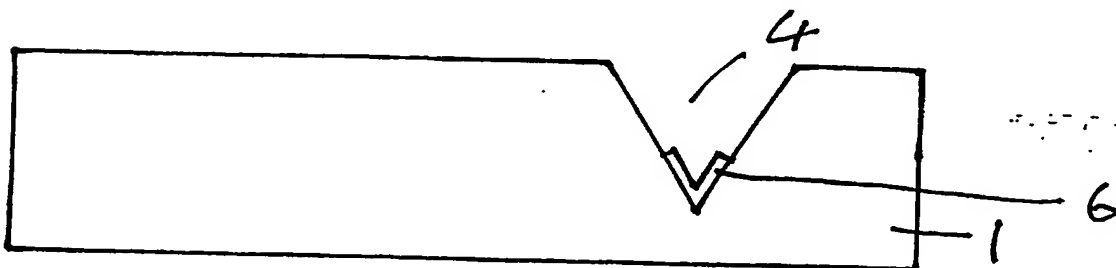
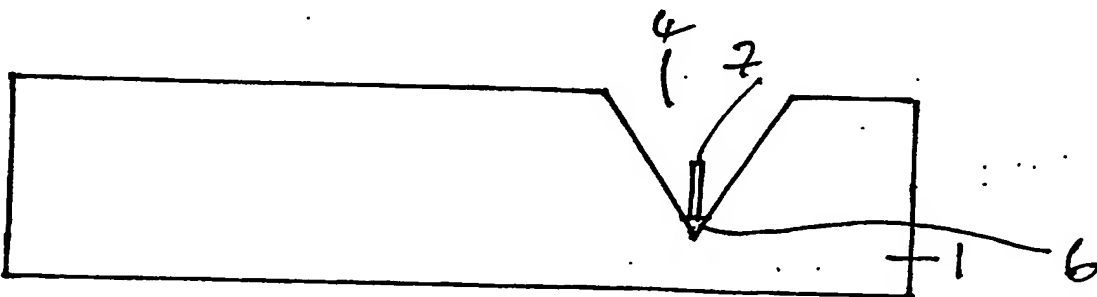


FIG. 1E



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2/6

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FIG. 1E

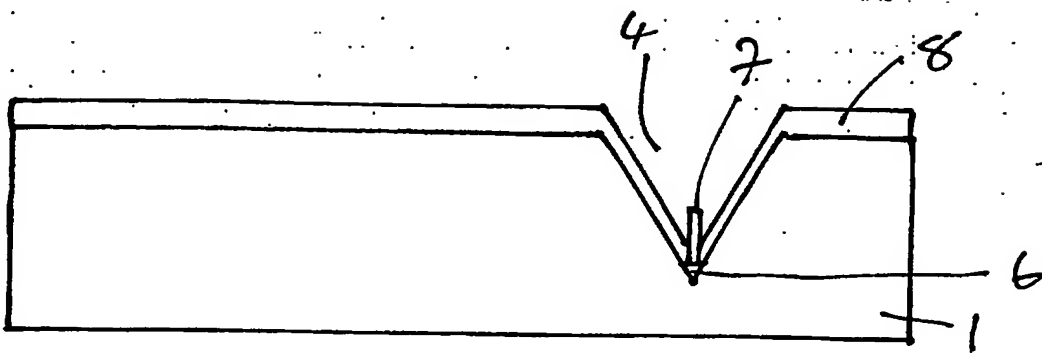


FIG. 1G

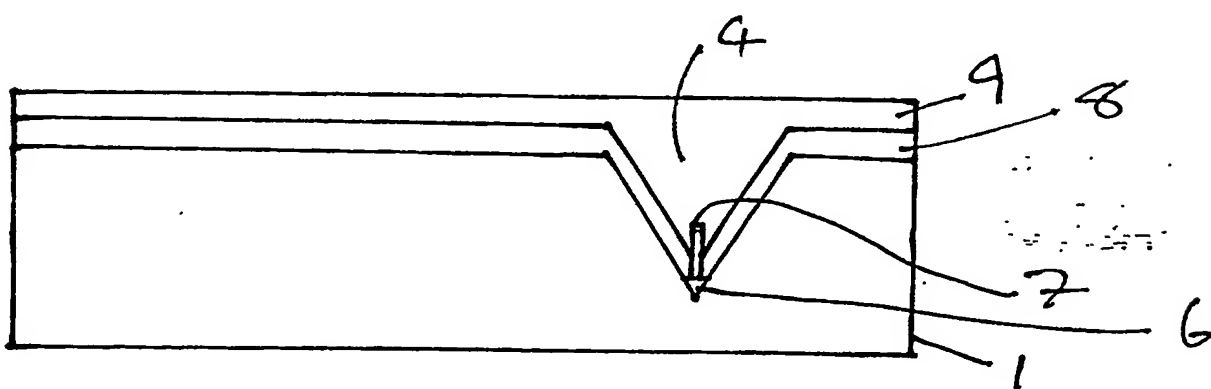


FIG. 1H

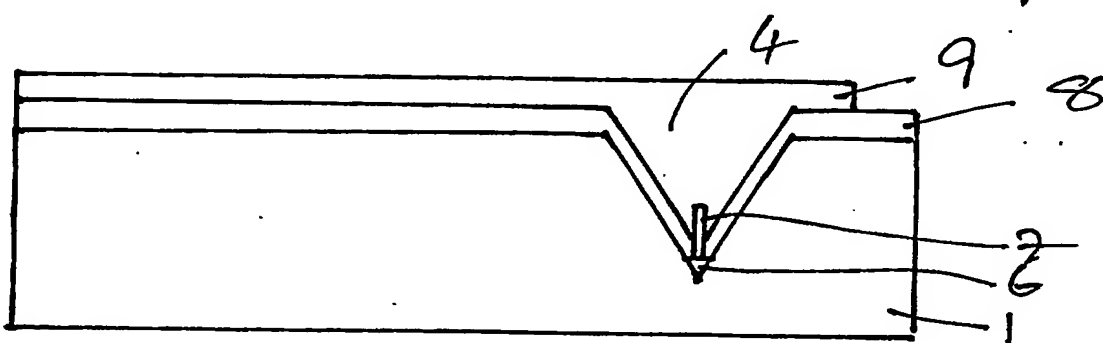
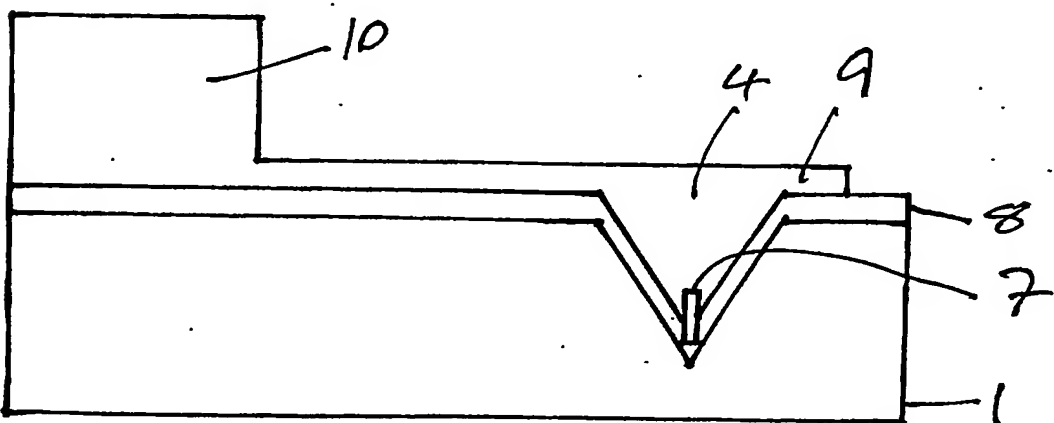


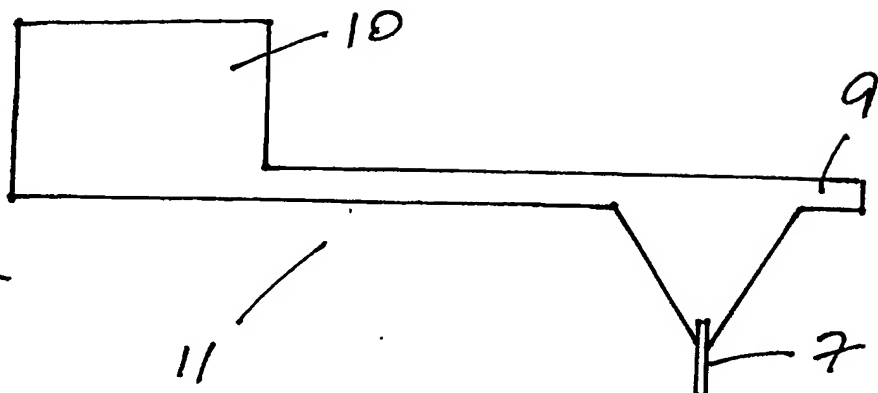
FIG. 1I



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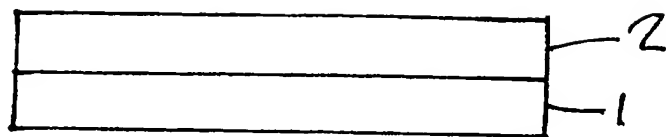
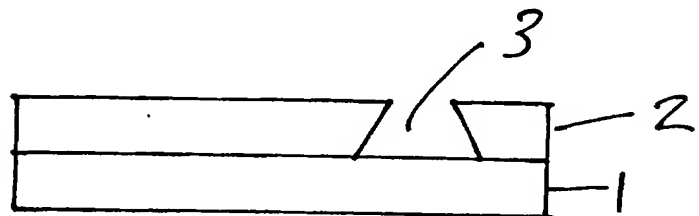
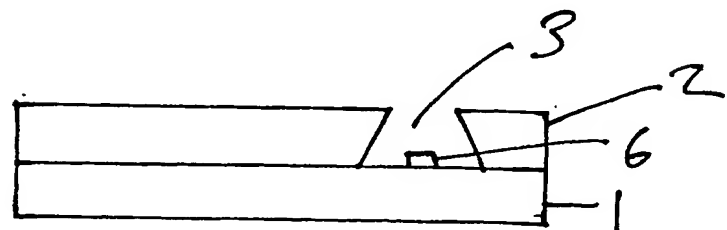
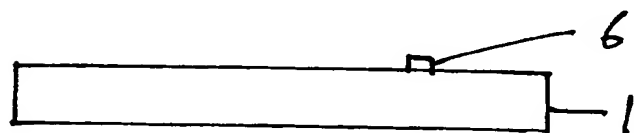
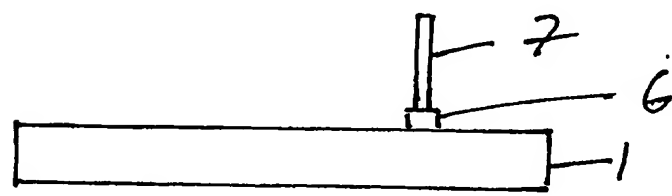
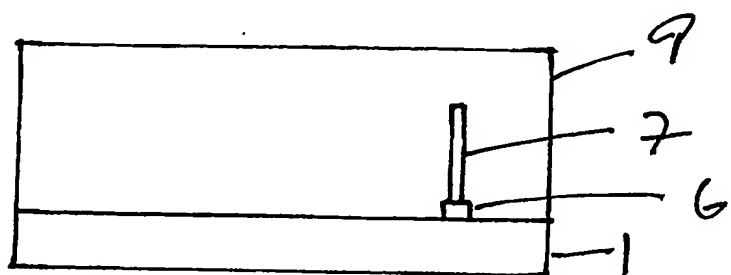
3/6

FIG. 1N



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FIG. 2AFIG. 2BFIG. 2CFIG. 2DFIG. 2EFIG. 2F

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5/6

FIG. 2G

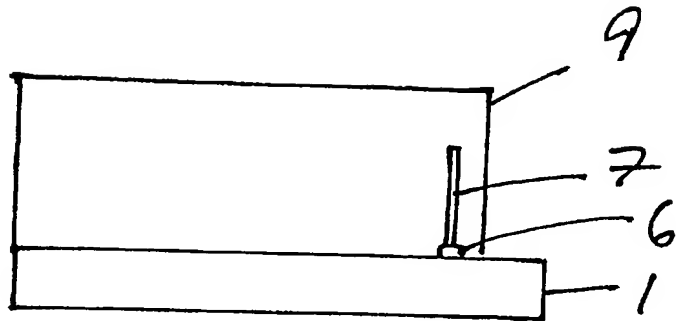


FIG. 2H

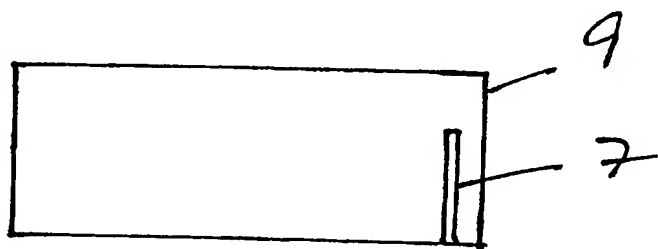
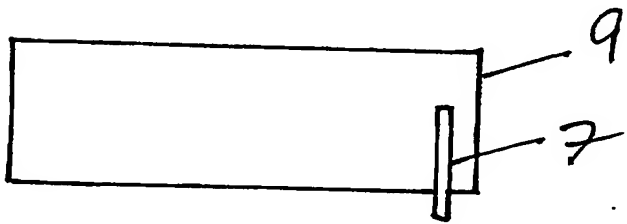


FIG. 2I



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6/6

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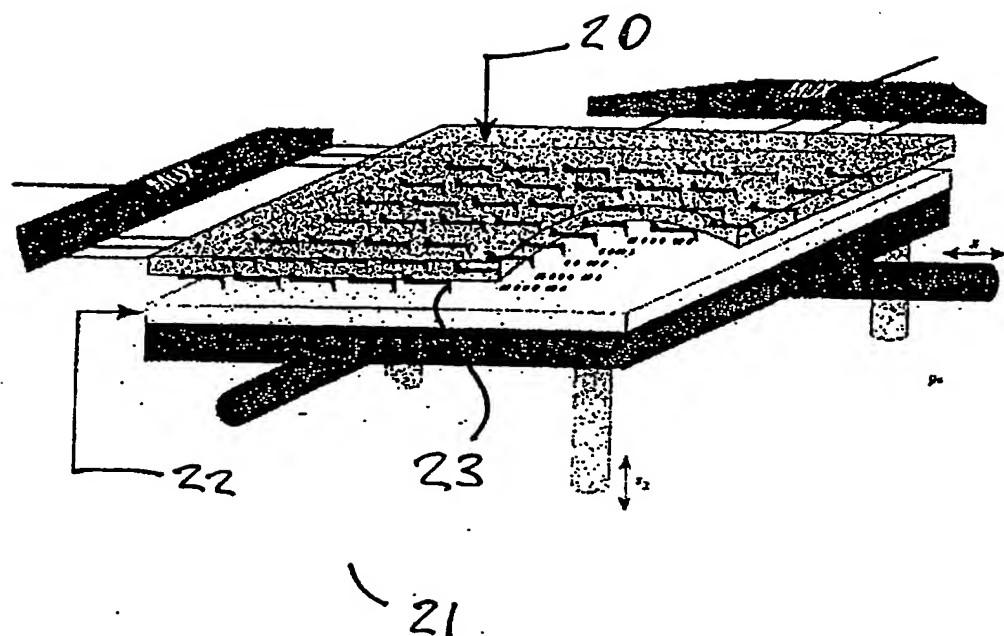


FIG. 8